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(54) Detergent composition having textile softening properties.

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DE - A - 2 646 995

GB - A - 1 400 898

US - A - 3 886 075

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Detergent composition having textile s ftening properties

Field of the invention

The present invention relates to detergent compositions which clean well and at the same time act as textile softeners.

δ Background of the invention

Numerous attempts have been made to formulate laundry detergent compositions which provide the good cleaning performance expected of them and which also have textile softening properties. Thus, attempts have been made to incorporate cationic textile softeners in anionic surfactant-based built detergent compositions employing various means of overcoming the natural antagonism between 10 the anionic and cationic surfactant species. For instance, in British patent specification 1,518,529, detergent compositions are described comprising organic surfactant, builders, and, in particulate form, a quaternary ammonium softener combined with a poorly water-soluble dispersion inhibitor which inhibits premature dispersion of the cationic in the wash liquor. Even in these compositions some compromise between cleaning and softening effectiveness has to be accepted. Another approach to 15 providing anionic detergent compositions with textile softening ability has been the use of smectitetype clays, as described in British patent specification 1,400,898. These compositions, although they clean well, require rather large contents of clay for effective softening, perhaps because the clay is not very efficiently deposited on the fabrics in the presence of anionic surfactants. Yet another approach to providing built detergent compositions with softening ability has been to employ nonionic surfactants zo instead of anionic with cationic softeners, and compositions of this type have been described in, for example, British patent specification 1,079,388, German Auslegeschrift 1,220,956 and US patent 3,607,763. However, it is found that if enough nonionic surfactant is employed to provide good cleaning, it impairs the softening effect of the cationic softener, so that, once again, a compromise between cleaning and softening effectiveness must be accepted.

The use of clay together with a water insoluble cationic compound and an electrically conductive metal salt as a softening composition adapted for use with anionic, nonionic, zwitterionic and amphoteric surfactants has been described in British patent specification 1,483,627.

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Recently it has been disclosed in German Offenlegungsschrift 2646995, that certain tertiary amines with two long chain alkyl or alkenyl groups and one short chain alkyl group are effective fabric softeners in detergent compositions when chosen to have an isoelectric point in the pH range such that they are in nonionic (amine) form in a normal alkaline wash liquor and are more in cationic (salt) form at the lower pH of a rinse liquor, and so become substantive to fabrics. Use of amines of this class, amongst others, in detergent compositions has also been previously disclosed in British patent specification 1,286,054.

Summary of the invention

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It has now been found that the combination of a certain class of tertiary amines and smectite-type clay in an alkaline detergent composition, or employed together with an alkaline detergent composition, provides pronounced textile softening benefits without imparing the cleaning performance of the detergent composition. Detergent compositions employing mixtures of clay with commercially available cationic textile softeners fail to provide the combination of softening and cleaning performance of the present compositions because of the incompatibility of the cationic softeners with alkaline detergent compositions, leading to an adverse effect upon their cleaning properties. In the present invention the softening effect is greater than that provided by the amine or the clay alone and the cleaning of particulate and soil stains is even enhanced.

According to the invention there is provided a textile softening detergent composition containing from 3% to 30% by weight of an organic surfactant, and from 10% to 80% of one or more water soluble inorganic or organic salts such that the pH of a 0.5% by weight aqueous solution of the composition is in the range from 8.5 to 11, wherein the textile softening component comprises the combination of (a) from 1% to 25% of a tertiary amine having the formula

wherein R_1 represents a C_{10} to C_{28} alkyl or alkenyl group, R_2 represents a group defined as for R_1 or a C_1 t C_7 alkyl group, and R_3 represents a C_1 to C_7 alkyl group, or where R_1 is a C_{16} — C_{28} alkyo group, R_2 may be a C_1 — C_7 alkyl group, r a mixtur of said amines, and

(b) from 1.5% to 35% of an impalpable smectite-type clay having an ion exchange capacity of at least 50 meq. per 100 grams.

It is preferr d that the weight ratio of t rtiary amin to clay b in th range fr m 10:1 to 1:10, preferably from 2:1 to 1:2. Pr f rably the pH f a 0.5% solution f the composition is in the range from 9.5 to 10.5.

5 Detailed description of the invention Organic surfactant

Anionic surfactant are much preferred for optimum combined cleaning and textile softening performance, but other classes of organic surfactants and mixtures thereof may be used, including surfactants such as the ethoxylated fatty alcohols and alkyl phenols well known in the art, amphoteric and zwitterionic surfactants and mixtures thereof as disclosed in US Patent No. 3,929,678, the disclosures of which are hereby incorporated by reference. When anionic surfactants are employed, it is preferred that nonionic and other classes of surfactant be absent but, if mixtures containing anionics are used, it is preferred that the anionic forms the major part of the mixture.

Suitable anionic non-soap surfactants are water soluble salts of alkyl benzene sulfonates, alkyl sulfates, alkyl polyethoxy ether sulfates, paraffin sulfonates, alphaolefin sulfonates, alphaolefin sulfonates, alkyl sulfonates, alkyl glyceryl ether sulfonates, fatty acid monoglyceride sulfates and sulfonates, alkyl phenol polyethoxy ether sulfates, 2-acyloxy-alkane-1-sulfonates, and beta-alkyloxy alkane sulfonates. Soaps are also suitable anionic surfactants.

Especially preferred alkyl benzene sulfonates have about 9 to about 15 carbon atoms in a linear or branched alkyl chain, more especially about 11 to about 13 carbon atoms. Sultable alkyl sulfates have about 10 to about 22 carbon atoms in the alkyl chain, more especially from about 12 to about 18 carbon atoms. Sultable alkyl polyethoxy ether sulfates have about 10 to about 18 carbon atoms in the alkyl chain and have an average of about 1 to about 12 —CH₂CH₂O— groups per molecule, especially about 10 to about 16 carbon atoms in the alkyl chain and an average of about 1 to about about 6—CH₂CH₂O— groups per molecule.

Suitable paraftin sulfonates are essentially linear and contain from about 8 to about 24 carbon atoms, more especially from about 14 to about 18 carbon atoms. Suitable alpha-olefin sulfonates have about 10 to about 24 carbon atoms, more especially about 14 to about 16 carbon atoms; alpha-olefin sulfonates can be made by reaction with sulfur trioxide followed by neutralization under conditions such that any sultones present are hydrolyzed to the corresponding hydroxy alkane sulfonates. Suitable alpha-sulfocarboxylates contain from about 6 to about 20 carbon atoms; included herein are not only the salts of alpha-sulfonated fatty acids but also their esters made from alcohols containing about 1 to about 14 carbon atoms.

Suitable alkyl glyceryl ether sulfates are ethers of alcohols having about 10 to about 18 carbon atoms, more especially those derived from coconut oil and tallow. Suitable alkyl phenol polyethoxy ether sulfates have about 8 to about 12 carbon atoms in the alkyl chain and an average of about 1 to about 6—CH₂CH₂O— groups per molecule. Suitable 2-acyloxy-alkane-1-sulfonates contain from about 2 to about 9 carbon atoms in the acyl group and about 9 to about 23 carbon atoms in the alkane moiety. Suitable beta-alkyloxy alkane sulfonates contain about 1 to about 3 carbon atoms in the alkyl group and about 8 to about 20 carbon atoms in the alkane moiety.

The alkyl chains of the foregoing non-soap anionic surfactants can be derived from natural sources such as coconut oil or tallow, or can be made synthetically as for example using the Ziegler or Oxo processes. Water solubility can be achieved by using alkali metal, ammonium, or alkanolammonium cations; sodium is preferred. Mixtures of anionic surfactants are contemplated by this invention; a satisfactory mixture contains alkyl benzene sulfonate having 11 to 13 carbon atoms in the alkyl group and alkyl sulfate having 12 to 18 carbon atoms in the alkyl group.

Suitable soaps contain about 8 to about 24 carbon atoms, more especially about 12 to about 18 carbon atoms. Soaps can be made by direct saponification of natural fats and oils such as coconut oil, tallow and fish oil, or by the neutralization of free fatty acids obtained from either natural or synthetic sources. The soap cation can be alkali metal, ammonium or alkanolammonium; sodium is preferred.

The compositions contain from 3 to 30% of organic detergent, preferably from 5 to 20% of anionic detergent.

The tertiary amines

Suitable amines are highly water insoluble amines of the structural formula

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wherein R_1 , R_2 and R_3 have the meanings define above. Preferably R_1 and R_2 each independently represents a C_{12} — C_{22} alkyl group, pr ferably straight chained, and R_3 is methyl, or ethyl. Suitabl 65 amines include

Di decyl methylamine

di lauryi methylamine

di myristyl methylamine

di cetyl methylamine

di stearyl methylamine

di arachadyl methylamine

di behenyl methylamine

arachadyl behenyl methylamine or

di (mixed arachidyi/behenyi) methylamine

di (tailowyl) methylamine

tallow dimethylamine

arachidyl/behenyl dimethylamine

and the corresponding ethyl amines, propylamines and butyl amines. Especially preferred is ditallowyl methylamine. This is commercially available as Kemamine T9701 (Humko Trade Name).

Other commercially available amines are Kemamine T1901 (DiC_{20/22} alkyi methylamine) and Kemamine T6501 (dicoconut methylamine).

The compositions contain from 1% to 25% usually from about 2% to about 15% by weight of the tertiary amine, especially from about 4% to about 8%.

20 The clay

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The smectite clays particularly useful in the practice of the present invention are sodium and calcium montmorillonites, sodium saponites, and sodium hectorites. The clays used herein have a particle size which cannot be perceived tactilely. Impalpable clays have particle sizes below about 50 microns; the clays used herein have a particle size range of from about 5 microns to about 50 microns.

The clay minerals can be described as expandable three-layer clays, i.e., alumino-silicates and magnesium silicates, having an ion exchange capacity of at least 50 meq/100 g. of clay and preferably at least 60 meq/100 g. of clay. The term "expandable" as used to describe clays relates to the ability of the layered clay structure to be swollen, or expanded, on contact with water. The three-layer expandable clays used herein are those materials classified geologically as smectites.

There are two distinct classes of smectite clays that can be broadly differentiated on the basis of the numbers of octahedral metal-oxygen arrangements in the central layer for a given number of silicon-oxygen atoms in the outer layers. The dioctahedral minerals are primarily trivalent metal ion-based clays and are comprised of the prototype pyrophyllite and the members mont-morillonite (OH)₄Si_{8-y}Al_y(Al_{4-x}Pe_x)O₂₀, and volchonskoite 35 (OH)₄Si_{8-y}Al_y(Al_{4-x}Cr_x)O₂₀, where x has a value of from 0 to about 4.0 and y has a value of from 0 to about 2.0. Of these only montmorillonites having exchange capacities greater than 50 meg/100 g. are suitable for the present invention and provide fabric softening benefits.

The trioctahedral minerals are primarily divalent metal ion based and comprise the prototype talc and the members hectorite (OH)₄Si_{8-y}Al_y(Mg_{8-x}Ll_x)O₂₀, saponite (OH)₄(Si_{8-y}Al_y)(Mg_{8-x}Al_x)O₂₀, sauconite 40 (OH)₄Si_{8-y}Al_y(Zn_{8-x}Al_x)O₂₀, vermiculite (OH)₄Si_{8-y}Al_y(Mg_{8-x}Fe_x)O₂₀, wherein y has a value of 0 to about 2.0 and x has a value of 0 to about 6.0. Hectorite and saponite are the only minerals in this class that are of value in the present invention, the fabric softening performance being related to the type of exchangeable cation as well as to the exchange capacity. It is to be recognized that the range of the water of hydration in the above formulas can vary with the processing to which the clay has been 45 subjected. This is immaterial to the use of the smectite clays in the present invention in that the expandable characteristics of the hydrated clays are dictated by the silicate lattice structure.

As noted hereinabove, the clays employed in the compositions of the instant invention contain cationic counterlons such as protons, sodium lons, potassium ions, calcium ions, and lithium ions. It is customary to distinguish between clays on the basis of one cation predominantly or exclusively absorbed. For example, a sodium clay is one in which the absorbed cation is predominantly sodium. Such absorbed cations can become involved in exchange reactions with cations present in aqueous solutions. A typical exchange reaction involving a smectite-type clay is expressed by the following equation:

Smectite clay (Na)+=smectite clay (NH₄)+NaOH

Since in the foregoing equilibrium reaction one equivalent weight of ammonium ion replaces an equivalent weight of sodium, it is customary to measure cation exchange capacity (sometimes termed "base exchange capacity") in terms of milli-equivalents per 100 g. of clay (meq/100 g). The cation exchange capacity of clays can be measured in several ways, including by electrodialysis, by exchange with ammonium ion followed by titration or by a methylene blue procedure, all as fully set forth in Grimshaw, "The Chemistry and Physics of Clays", pp. 264—265, Interscience (1971). The cation exchange capacity of a clay mineral relat s to such factors as the expandable properties of the clay, the charg of the clay, which, in turn, is determined at least in part by the lattice structure and the like. The ion exchange capacity of clays varies widely in the range from about 2 meq/100 g, for kaolinites to

about 150 meg/100 g., and greater, for certain smectite clays. Illite clays although having a three layer structure, are f an n-expanding lattice type and have an ion xchange capacity somewhere in the lower portion of the range, i.e., around 26 meg/100 g. for an average illite clay. Attapulgites, another class of clay minerals, have a spicular (i.e. needle-like) crystalline form with a low cation xchange capacity (25—30 meg/100 g.). Their structure is composed of chains of silica tetrahedrons linked together by octahedral groups of oxygens and hydroxyls containing AI and Mg atoms.

It has been determined that illite, attapulgite, and kaolinite clays, with their relatively low ion exchange capacities, are not useful in the present compositions. However the alkali metal montmorillonites, saponites, and hectorites, and certain alkaline earth metal varieties of these minerals such as to calcium montmorillonites have been found to show useful fabric softening benefits when incorporated

in the compositions in accordance with the present invention.

Specific non-limiting examples of such fabric softening smectite clay minerals are:

Sodium Montmorillonite

Brock
Volclay BC
Gelwhite GP
Thixo-Jel #
Ben-A-Gel

Sodium Hectorite
Veegum F
Laponite SP

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25 Sodium Saponite Barasym NAS 100

Calcium Montmorillonite Soft Clark 30 Gelwhite L Imvite K

> Lithium Hectorite Barasym LIH 200

Accordingly, smectite clays useful herein can be characterised as montmorillonite, hectorites, and saponite clay minerals having an ion exchange capacity of at least about 50 meg/100 g. and preferably at least 60 meg/100 g. Most of the smectite clays useful in the compositions herein are commercially available under various trade names for example Thixogel No. 1 and Gelwhite GP from Georgia. Kaolin Co., Elizabeth, New Jersey; Imvite K from Industrial Mineral Ventures; Volclay BC and Volclay # 325, from America Colloid Co., Skokie, Illinois; and Veegum F, from R. T. Vanderbilt. It is to be recognised that such smectite minerals obtained under the foregoing tradenames can comprise mixtures of the various discrete mineral entities. Such mixtures of the smectite minerals are suitable for use herein.

Within the classes of montmorillonite, hectorite, and saponite clay minerals having a cation exchange capacity of at least about 50 meg/100 g, certain clays are preferred for fabric softening purposes. For example, Gelwhite GP is an extremely white form of smectite clay and is therefore preferred when formulation white granular detergent compositions. Volclay BC, which is a smectite clay mineral containing at least 3% of iron (expressed as Fe₂O₃) in the crystal lattice, and which has a very high ion exchange capacity, is one of the most efficient and effective clays for use in detergent softening compositions. Invite K is also very satisfactory.

Appropriate clay minerals for use herein can be selected by virtue of the fact that smectites exhibit a true 14 Å x-ray diffraction pattern. This characteristic pattern, taken in combination with exchange capacity measurements performed in the manner noted above, provides a basis for selecting particular smectite-type minerals for use in the compositions disclosed herein.

The smectite clay materials useful in the present invention are hydrophilic in nature, i.e. they display swelling characteristics in aqueous media. Conversely they do not swell in nonaqueous or predominantly nonaqueous systems.

The compositions contain from 1.5% to 35%, preferably from about 4% to about 15% of said smectite-type clay, especially from about 5—12%.

60 Water-soluble salts

The compositions of the invention contain from 10% to 80% of water soluble salts, preferably from 20% to 70%, and most usually from 30% to 60%, and these may be any which are such that the detergent c mposition in a 0.5% by weight aqueous solution has pH in the specified range, that is from 8.5 to 11, preferably from 9.5 to 10.5. At this pH the tertiary amines of the inv ntion are in nonionic (amine) form and are therefore compatible with an inic surfactants.

Preferably the water soluble salts are detergency build rs and th se can b of the polyvalent inorganic and polyvalent organic types, or mixtures thereof. Non-limiting examples of suitable water-solubl inorganic alkaline det rgent builder salts include th alkali metal carbonates, borates, phosphat s, polyphosphates, tripolyphosphates, bicarbonates, and silicates. Specific examples of such salts include the sodium and potassium tetraborates, bicarbonates, carbonates, tripolyphosphates, pyrophosphates, pentapolyphosphates, and hexametaphosphates. Sulphates are usually also present.

Examples of suitable organic alkaline detergency bullders salts are:
(1) water-soluble amino polyacetates, e.g., sodium and potassium ethylenediaminetetraacetates, nitrilotriacetates, N-(2-hydroxyethyl) nitrilodiacetates and diethylenetriamine penta-

10 acetates;

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(2) water-soluble salts of phytic acid, e.g. sodlum and potassium phytates;

(3) water-soluble polyphosphonates, including sodium, potassium and lithium salts of methylenediphosphonic acid and the like and aminopolymethylene phosphonates such as ethyldiaminetetramethylenephosphonate and diethylenetriaminepentamethylene phosphonate, and polyphosphonates as described in German Application DOS 2816770.

(4) water-soluble polycarboxylates such as the salts of lactic acid, succinic acid, malenic acid, carboxymethylsuccinic acid, 2-oxa-1,1,3-propane tricarboxylic acid, 1,1,2,2-ethane tetracarboxylic acid, cyclopentane-cis, cis, cis-tetracarboxylic acid, mellitic

acld and pyromellitic acid.

Mixtures of organic and/or inorganic builders can be used herein. One such mixture of builders is disclosed in Canadian Patent No. 755,038, e.g. a ternary mixture of sodium tripolyphosphate, tri-

sodium nitrilotriacetate, and trisodium ethane-1-hydroxy-1,1-diphosphonate.

Another type of detergency builder material useful in the present compositions and processes comprises a water-soluble material capable of forming a water-insoluble reaction production with water hardness cations preferably in combination with a crystallization seed which is capable of providing growth sites for said reaction product. Such "seeded builder" compositions are fully disclosed in British Patent Specification No. 1,424,406.

Preferred water soluble bullders are sodium tripolyphosphate and sodium silicate, and usually both are present. In particular it is preferred that a substantial proportion, for instance from 3 to 15% by weight of the composition of sodium silicate (solids) of ratio (weight ratio SiO₂:Na₂O) from 1:1 to 3.5:1

be employed.

A further class of detergency builder materials useful in the present invention are insoluble sodium alumino-silicates, particularly those described in Belgian Patent 814,874, issued November 12, 1974 incorporated herein by reference. This patent discloses and claims detergent compositions containing sodium aluminosilicates of the formula

$Na_z(AIO_2)_z(SiO_2)_y x H_2O$

wherein Z and Y are integers equal to at least 6, the molar ratio of Z to Y is in the range of from 1.0:1 to about 0.5:1 and x is an integer from about 15 to about 264. A preferred material is $Na_{12}(SiO_2AIO_2)_{12}27H_2O$.

Preferably, the compositions contain from 20% to 70% of builders, more usually 30% to 60% by weight. If present, incorporation of about 5% to about 25% by weight of aluminosilicate is suitable, partially replacing water soluble builder salts, provided that sufficient water soluble alkaline salts remain to provide the specified pH of the composition in aqueous solution.

Optional components

The optional components usual in built laundry detergents may of course be present. These include bleaching agents such as sodium perborate, sodium percarbonate and other perhydrates, at levels from about 5% to 35% by weight of the composition, and activators therefor, such as tetra acetyl ethylene diamine, tetra acetyl glycouril and others known in the art, and stabilisers therefor, such as magnesium silicate, and ethylene diamine tetra acetate.

Suds controlling agents are often present. These include suds boosting or suds stabilising agents such as mono- or di-ethanolamides of fatty acids. More often in modern detergent compositions, suds, suppressing agents are required. Soaps especially those having 16—22 carbon atoms, or the corresponding fatty acids, can act as effective suds suppressors if included in the anionic surfactant component of the present compositions. Usually about 1% to about 4% of such soap is effective as a suds suppressor. Very suitable soaps when suds suppression is a primary reason for their use, are those derived from Hyfac (Trade name for hardened marine oil fatty acids predominantly C₁₈ to C₂₀).

However, non-soap suds suppressors are preferred in synthetic detergent based compositions of the invention since soap or fatty acid tends to give rise to a characteristic odour in these compositions.

Preferred suds suppressors comprise silicones. In particular there may be employ dia particulate suds suppressor comprising silicone and silanated silica releasably inclosed in water soluble or dispersible substantially non-surfac active detergent impermeable carrier. Suds suppressing agent of this sort are disclosed in British patent specification 1,407,997. A very suitable granular (prilled) suds

suppressing product comprises 7% silica/silicone (85% by weight silanated silica, 15% silicone, obtained from Messrs. Dow Corning), 65% sodium tripolyphosphate, 25% Tallow alcohol condensed with 25 molar prop rtions of ethylene oxid, and 3% m isture. The amount of silica/silic ne suds suppressor empl yed depends upon the degree f suds suppression desired but is fiten in the range 5 from 0.01% to 0.5% by weight of the detergent composition. Oth r suds suppressors which may be used are water insoluble, preferably microcrystalline, waxes having melting point in the range from 35 to 125°C and saponification value less than 100, as described in British patent specification

1,492,938.
Yet other suitable suds suppressing systems are mixtures of hydrocarbon oil, a hydrocarbon oil, a hydrocarbon oil and, suitable suds suppressing systems are mixtures of hydrocarbon oil, a hydrocarbon oil and, suitable suds suppressing systems are mixtures of hydrocarbon oil, a hydrocarbon oil and, successing systems are mixtures of hydrocarbon oil, a hydrocarbon oil and successing systems are mixtures of hydrocarbon oil and succession 10 wax and hydrophobic silica as described in European laid open patent application No. 0000216 and, especially, particulate suds suppressing compositions comprising such mixtures, combined with a nonionic ethoxylate having hydrophilic lipophilic balance in the range from 14—19 and a compatibilising agent capable of forming inclusion compounds, such as urea. These particulate suds suppressing compositions are described in European patent application 79200472.3 filed 29 August,

Soil suspending agents are usually present at about 0.1 to 10%, such as water soluble salts of carboxymethylcellulose, carboxyhydroxymethyl cellulose, polyethylene glycols of molecular weight from about 400 to 10000 and copolymers of methylvinylether and maleic anhydride or acid, available from the General Aniline and Film Corporation under the Trade Name Gantrez.

Proteolytic, amylolytic or lipolytic enzymes, especially proteolytic, and optical brighteners, of anionic cationic or nonlonic types, especially the derivatives of sulphonated triazinyl diamino stilbene may be present. A further useful additive is a photo activated bleach comprising a mixture of the tri and tetra sulphonated derivatives of zinc phthalocyanine as described in B.P. Specification Nos. 1372035 and 1408144.

Through the description herein, where sodium salts have been referred to, potassium, lithium or ammonium or amine salts may be used instead if their extra cost etc. are justified for special reasons,

Preparation of the compositions

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The detergent compositions may be prepared in any way, as appropriate to their physical form, 30, as by mixing the components, co-agglomerating them or dispersing them in a liquid carrier. Preferably the compositions are granular and are prepared by spray drying an aqueous slurry of the non-heatsensitive components to form spray dried granules into which may be admixed the heat sensitive components such as persalts, enzymes, perfumes etc. Although the amine may be included in the slurry for spray drying, it is preferred that it be incorporated by being sprayed in liquid form on the spray dried 35 granules before or after other heat sensitive solids have been dry mixed with them. Although the amine is generally a waxy solid of rather low melting point the granules so made are surprisingly crisp and free-flowing. Alternatively the amine in liquid form may be sprayed onto any particulate component or components of the composition which are able to act as carrier granules. The clay component may be added to the slurry for spray drying or may be dry mixed, as preferred for reasons unrelated to its softening effect, such as for optimum colour of the product.

Examples 1 and 2 Textile softening detergent compositions were prepared having the formula, in parts per cent by

45	Example	1	2
	(a) Sodium linear dodecylbenzene sulphonate (LAS)	8	8
	(a) Sodium tripolyphosphate	32	30
	(a) Sodium silicate (ratio SiO ₂ /NaO2)	6	6
50	(a) Sodium sulphate	5	5
	(c) Sodium perborate	25	22
	(a) Sodium carboxymethyl cellulose	0.8	0.8
	(a) Sodium ethylenediamine tetra acetate	0.2	0.2
	(c) Enzyme granules	0.4	0.4
55	(a) Optical brightener	0.2	0.2
	(b) Perfume	0.25	0.25
	(c) Silica-silicone suds suppressor*	0.15	0.15
	(a) Clay** (montmorillonite)	10	10
	(b) Ditallowyl methylamine	6	12
60	— Moisture etc.	6	5

^{*}Silica-polydimethyl siloxane in ratio by weight 90:10

The compositions were prepared by making spray dried granules containing comp nents (a), 65 spraying m Iten ditall wylmethylamin and perfume (comp nents (b)) on t th m in a r tating drum,

^{**&}quot;Imvite K—Tradenam of Messrs. Industrial Mineral Ventures (I.M.V.).

and dry mixing the resultant granules with components (c). 0.5% solutions of the compositions in water at 20°C had pH 8.9 to 10.1.

These comp sitions had as good cleaning performance as the same compositions lacking the clay and amine, with slightly better cleaning performanc on clay soiling. Cotton test pieces washed with these compositions were softer in feel than similar test pi ces washed with the same detergent compositions excluding either the amine or the clay or both.

Further it was found that the softening effect provided by the clay was greater when the clay was added to the amine containing detergent composition of Example 1 than when it was added to the detergent composition of Example 1 lacking amine.

Similar performance is obtained when the tertiary amine is replaced by dicoconut methylamine. di-myristyl methylamine, ditallowyl ethylamine, di(arachidyl behenyl) methylamine, ditallowyl propylamine, or tallow dimethylamine.

Similar performance is obtained when the "Imvite K" clay is replaced by Volclay BC, Gelwhite GP, Soft Clark, or Gelwhite L. Volclay is a tradename of American Colloids Co., Gelwhite and Soft Clark are Tradenames of Georgia Kaolin Co.

Similar performance is obtained when the LAS is replaced by a mixture of 4% LAS and 4% sodium coconut alkyl sulphate, or a mixture of 5% LAS and 3% sodium tallow alkyl sulphate.

Similar performance was obtained when the clay was dry mixed together with components (c) instead of being added to the slurry for spray drying.

Examples 3 to 7

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The following compositions are prepared substantially as described in Example 1, and provide cleaning and textile softening benefits. Quantities are in parts per cent by weight.

25	Example	3	4	5	6	7	
	Sodium linear dodecyl benzene sulphonate	15	5	8	10	_	-
	Sodium tallow alkyl sulphate	_	5	_	_		
	Sodium soap (80/20 Tallow-coconut)		3		_	45	
3 0	Sodium tripolyphosphate	30	44	12	5	5	
	Sodium carbonate	4		_	14	20	
	Sodium silicate	8	6	10	8	10	
	Sodium sulphate	12	8	6	8		
	Sodium perborate tetrahydrate	7	10	20	_		
35	Sodium alumino silicate	-		20	_		
	Sodium carboxymethyl cellulose	1	1	1	1		
	Sodium ethylenediamine tetra acetate	0.2	0.2	0.2	·		
	Enzyme granules	0.5	0.5	0.5		_	
	Optical brightener	0.3	0.3	0.3		0.3	
40	Clay (Imvite K)	4	8 8		30	-	
		-	_	10	30	3	
	Ditallow methylamine	10	2	6	20	4	
	Moisture etc.	8	7	6	4	12.7	

Claims

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1. A textile softening detergent composition containing from 3% to 30% by weight of an organic surfactant, and from 10% to 80% of one or more water soluble inorganic or organic salts such that the pH of a 0.5% by weight aqueous solution of the composition is in the range from 8.5 to 11 characterised in that it comprises as the textile softening component, the combination of

(a) from 1% to 25% of a tertiary amine having the formula

wherein R₁ represents a C₁₀ to C₂₈ alkyl or alkenyl group, R₂ represents a group defined as for R₁ or where R₁ is a C₁₆—C₂₈ alkyl group, R₂ may be a C₁ to C₇ alkyl group, and R₃ represents a C₁ to C₇ alkyl group, or of a mixture of said amines, and
(b) from 1.5% to 35% of an impalpable smectite-type clay having an ion exchange capacity of at

least 50 meq. per 100 grams.

2. A detergent compositi n according to Claim 1 which contains from 5% to 20% f an anionic surfactant selected from s dium or potassium C_{9-15} alkyl benzene sulphonates and C_{12-18} alkyl 65 sulphates and mixtures thereof.

3. A deterg nt composition according to Claims 1 and 2 wherein the weight ratio of tertiary amine to smectit -type clay is in the range from 10:1 to 1:10, preferably from 2:1 t 1:2.

4. A detergent c mpositi n according t any on of Claims 1 to 3 which contains by weight from 2% to 15% of an amine of formula wherein R₁ and R₂ each indep indently represents a C₁₂ to C₂₂ alkyl group and R₃ Is methyl r thyl.

5. A detergent composition according to Claim 4 wherein the amine is ditallowyl methylamine. 6. A detergent composition according to any one of Claims 1 to 4 which contains by weight from 4% to 15% of a smectite-type clay selected from the group consisting of alkali metal and alkali earth metal montmorillonites, saponites, hectorites or mixtures thereof.

A detergent composition according to any one of the preceding claims which contain from 20% to 70% of the water soluble salts, which salts comprise detergency builders selected from alkaline sodium and potassium carbonates, borates, phosphates, polyphosphates, silicates, polycarboxylates, polyphosphonates and aminopolycarboxylates.

8. A detergent composition according to Claim 6 also containing from about 5% to about 25% by weight of the composition of a water insoluble aluminosilicate of formula Na_z(AlO_z)_z(SlO_z)_zxH_zO wherein z and y are integers equal to at least 6, the molar ratio of z to y is in the range from 1:1 to 0.5:1 and x is an integer from about 15 to about 264.

9. A detergent compositions according to any one of the preceding claims which also contains a suds suppressor selected from C₁₈ to C₂₄ fatty acids or soaps, microcrystalline waxes, silicone-hydro-20 phobic silica mixtures, combinations of paraffin oil, wax and hydrophobic silica, and mixtures thereof.

10. A method of preparing a detergent composition according to any one of Claims 1 to 9, wherein the amine, in liquid form, is sprayed on to preformed granules comprising some or all of the other components of the composition.

25 Patentansprüche

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1. Textilweichmacher-Waschmittelzusammensetzung enthaltend 3—30 Gew.% eines organischen Tensides und 10-80 % eines wasserlöslichen anorganischen oder organischen Salzes oder mehrerer solcher Salze derart, dass der pH-Wert einer 0,5 gewichtsprozentigen wässrigen Lösung der 30. Zusammensetzung im Bereich von 8,5-11 liegt, dadurch gekennzeichnet, dass sie als Textilweichmacherkomponente die Kombination aus

(a) 1-25 % eines tertiären Amins der Formel

in der R₁ eine C₁₀—C₂₈-Alkyl- oder -Alkenylgruppe, R₂ eine wie für R₁ definierte Gruppe darstellt oder, wenn R
1 eine C₁₆—C₂₆-Alkylgruppe ist, R
2 eine C₁—C₇-Alkylgruppe sein kann, und R
3 eine C₁—C₇-Alkylgruppe darstellt, oder einer Mischung solcher Amine, und 1,5—35 % eines Charles baren smektitartigen Tons mit einer lonenaustauschkapazität

von mindestens 50 Milliäquivalent pro 100 Gramm enthält.

2. Waschmittelzusammensetzung nach Anspruch 1, die 5-20 % eines anionischen Tensides gewählt aus Natrium- oder Kalium-C9-15-alkylbenzolsulfonaten und C12-18-Alkylsulfaten sowie Mischungen hiervon enthält.

3. Waschmittelzusammensetzung nach Anspruch 1 und 2, bei welcher das Gewichtsverhältnis von tertiärem Amin zum smektitartigen Ton im Bereich von 10:1 bis 1:10, vorzugsweise von 2:1 bis

4. Waschmittelzusammensetzung nach einem der Ansprüche 1-3, die, bezogen auf das Gewicht, 2—15 % eines Amins der Formel enthält, in der R₁ und R₂ jeweils unabhängig C₁₂—C₂₂-Alkylgruppen darstellen und R₃ Methyl oder Ethyl ist.

5. Waschmittelzusammensetzung nach Anspruch 4, in der das Amin Di-talgalkyl-methylamin ist. 6. Waschmittelzusammensetzung nach einem der Ansprüche 1-4, die, bezogen auf das Gewicht, 4-15 % eines smektitartigen Tons gewählt aus der Gruppe bestehend aus Alkalimetall- und Erdalkalimetall-montmorilloniten, -saponiten, -hectoriten oder Mischungen hiervon enthält.

7. Waschmittelzusammensetzung nach einem der vorangehenden Ansprüche, die 20—70 % der wasserlöslichen Salze enthält, welche Salze Detergens-Gerüststoffe gewählt aus alkalischen Natriumund Kaliumcarbonaten, -boraten, -phosphaten, -polyphosphaten, -silicaten, -polycarboxylaten, -polyphosphonaten und -aminopolycarboxylaten enthalten.

8. Waschmitt Izusammensetzung nach Anspruch 6, die etwa 5 bis etwa 25 %, bezogen auf das Gewicht der Zusammensetzung, eines wass runi slich n Aluminlumsilicat s d r Formel Na_z(AlO₂)_z-(SiO₂)_xH₂O enthält, in der z und y ganz Zahlen gleich mindest ns 6 sind, das Molverhältnis v n z zu y 65 im Bereich v n 1:1 bis 0,5:1 liegt und x ine ganze Zahl von twa 15 bis twa 264 ist.

9. Waschmittelzusammensetzung nach einem der v rangehenden Ansprüche, die ausserdem ein Schaumunterdrückungsmittel gewählt aus C_{16} — C_{24} -Fettsäuren der -seif n, mikr kristallinen Wachsen, sllicon-hydrophoben Sillciumdloxidmischungen, Kombinationen aus Paraffin i, Wachs und hydrophobem Sillciumdloxid und Mischungen hiervon enthält.

10. Verfahren zur Herstellung einer Waschmittelzubereitung nach einem der Ansprüche 1—9, bei welchem das Amin in flüssiger Form auf vorgeformte K\u00f6rner gespr\u00fcht wird, die einige oder alle Komponenten der Zusammensetzung enthalten.

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Revendications

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1. Une composition détergente adoucissante sur textiles contenant 3 % à 30 % en poids d'un tensio-actif ou agent de surface organique et 10 % à 80 % d'un ou de plusieurs sels organiques ou minéraux solubles dans l'eau tels que le pH d'une solution aqueuse à 0,5 % en poids de a composition soit compris entre 8,5 et 11, caractérisée en ce qu'elle comprend, en tant que composant adoucissant des textiles, la combinaison

(a) de 1 % à 25 % d'une amine tertiaire répondant à la formule:

R₁N—R₃

dans laquelle R₁ représente un groupe alkyle ou alcényle en C₁₀ à C₂₆, R₂ représente un groupe défini comme R₁ ou, lorsque R₁ est un groupe alkyle en C₁₈—C₂₆, R₂ peut être un groupe alkyle en C₁ à C₇ et R₃ représente un groupe alkyle en C₁ à C₇ ou d'un mélange desdites amines, et (b) de 1,5 % à 35 % d'une argie implaable de type smectite ayant une capacité d'échange

d'ions d'au moins 50 méq. pour 100 g.

Composition détergente selon la revendication 1, caractérisée en ce qu'elle contient 5 % à 20 % d'un agent de surface anionique choisi parmi les alkyl C₉₋₁₅ benzène sulfonates et alkyl C₁₂₋₁₈ sulfates de sodium ou de potassium et leurs mélanges.

3. Composition détergente selon les revendications 1 et 2, caractérisée en ce que le rapport pondéral de l'amine tertiaire à l'argile de type smectite se situe dans la gamme de 10:1 à 1:10, de préférence de 2:1 à 1:2.

4. Composition détergente selon l'une quelconque des revendications 1 à 3, caractérisée en ce qu'elle contient en poids 2 % à 15 % d'une amine répondant à la formule dans laquelle R_1 et R_2 représentent chacun indépendamment un groupe aikyle en C_{12} à C_{22} et R_3 est un méthyle ou un éthyle.

5. Composition détergente selon la revendication 4, caractérisée en ce que l'amine est la di-suif méthylamine.

6. Composition détergente selon l'une quelconque des revendications 1 à 4, caractérisée en ce qu'elle contient en polds 4 % à 15 % d'une argile de type smectite choisie dans le groupe consistant en montmorillonites, saponites, hectorites de métaux alcalins et de métaux alcalino-terreux ou en leurs mélanges.

7. Composition détergente selon l'une quelconque des revendications précédentes, caractérisée en ce qu'elle contient 20 % à 70 % des sels solubles dans l'eau, lesdits sels comprenant des adjuvants de détergence choisis parmi les carbonates, borates, phosphates, polyphosphates, silicates, polycarboxylates, polyphosphonates et aminopolycarboxylates de sodium et de potessium alcalins.

8. Composition détergente selon la revendication 6, caractérisée en ce qu'elle contient également environ 5 % à environ 25 % en poids par rapport à la composition d'un aluminosilicate insoluble dans l'eau de formule Na_x(AlO₂)_x(SlO₂)_xxH₂O dans laquelle z et y sont des nombres entiers égaux à au moins 6, le rapport molaire de z à y se situe entre 1:1 et 0,5:1 et x est un nombre entier d'environ 15 à environ 264.

9. Composition détergente selon l'une quelconque des revendications précédentes, caractérisée en ce qu'elle contient également un suppresseur de mousse choisi parmi des acides gras ou des savons en C₁₈ à C₂₄, des cires microcristallines, des mélanges de silicone-silice hydrophobe, des combinaisons d'huile de paraffine, de cire et de silice hydrophobe et leurs mélanges.

10. Procédé de préparation d'une composition détergente selon l'une quelconque des revendications 1 à 9, caractérisée en ce que l'amine est pulvérisée sous forme liquide sur des granules préformés contenant une partie ou la totalité des autre constituants de la composition.

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